

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

**(19) World Intellectual Property Organization
International Bureau**



A standard linear barcode is located at the bottom of the page, spanning most of the width. It is used for document tracking and identification.

(43) International Publication Date
17 October 2002 (17.10.2002)

PCT

(10) International Publication Number
WO 02/081721 A2

(51) International Patent Classification⁷: C12P 13/00

GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW,
MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG,
SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN,
YU, ZA, ZM, ZW.

(21) International Application Number: PCT/EP02/02419

(22) International Filing Date: 6 March 2002 (06.03.2002)

(25) Filing Language:

English

(84) **Designated States (regional):** ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

(26) Publication Lang

English

(30) Priority Data:

101 16 518.8 3 April 2001 (03.04.2001) DE

(71) **Applicant:** DEGUSSA AG [DE/DE]; Bennigsenplatz 1,
40474 Düsseldorf (DE).

Published:

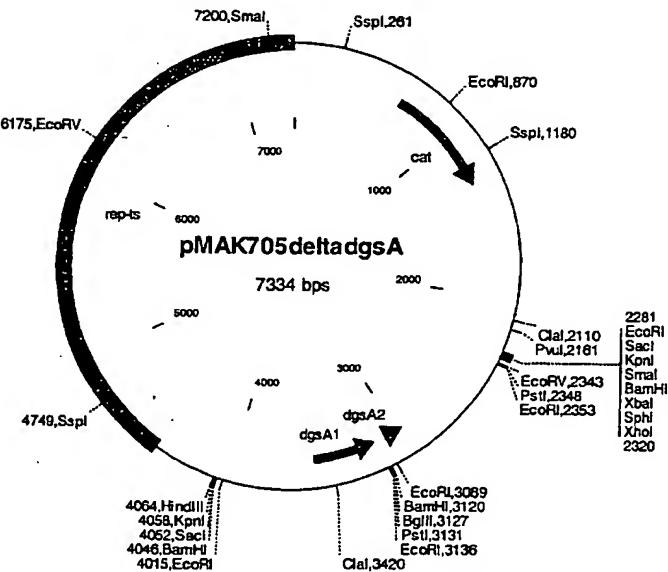
— without international search report and to be republished upon receipt of that report

- with (an) indication(s) in relation to deposited biological material furnished under Rule 13bis separately from the description

(72) **Inventors:** RIEPING, Mechthild; Mönkebergstrasse 1, 33619 Bielefeld (DE). HERMANN, Thomas; Zirkonstrasse 8, 33739 Bielefeld (DE).

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: PROCESS FOR THE PRODUCTION OF L-AMINO ACIDS USING STRAINS OF THE FAMILY ENTEROBACTERIACEAE THAT CONTAIN AN ATTENUATED DGSA GENE



W302/081721 A2

WO 2010/043330
(57) Abstract: The invention relates to a process for the production of L-amino acids, in particular L-threonine, in which the following steps are carried out: a) attenuation of the dgsA gene or nucleotide sequences coding therefor for L-threonine producing the "design" L-amino acid, in which the dgsA gene or nucleotide sequences coding therefor are attenuated, in particular are switched off, b) enrichment of the L-amino acid in the medium or in the cells of the bacteria, and c) isolation of the L-amino acid.

**Process for the Production of L-Amino Acids Using
Strains of the Family Enterobacteriaceae
that Contain an Attenuated dgsA Gene**

Field of the Invention

5 The present invention relates to a process for the enzymatic production of L-amino acids, in particular L-threonine, using strains of the family Enterobacteriaceae in which the dgsA gene is attenuated.

Prior Art

10 L-amino acids, in particular L-threonine, are used in human medicine and in the pharmaceutical industry, in the foodstuffs industry, and most especially in animal nutrition.

It is known to produce L-amino acids by fermentation of
15 strains of Enterobacteriaceae, in particular Escherichia coli (E. coli) and Serratia marcescens. On account of their great importance efforts are constantly being made to improve processes for producing the latter. Process improvements may relate to fermentation technology
20 measures, such as for example stirring and provision of oxygen, or the composition of the nutrient media, such as for example the sugar concentration during the fermentation, or the working-up to the product form, for example by ion exchange chromatography, or the intrinsic
25 performance properties of the microorganism itself.

Methods comprising mutagenesis, selection and mutant choice are employed in order to improve the performance properties of these microorganisms. In this way strains are obtained that are resistant to antimetabolites, such as for example
30 the threonine analogue α -amino- β -hydroxyvaleric acid (AHV) or are auxotrophic for regulatorily important metabolites,

and that produce L-amino acids such as for example L-threonine.

Methods of recombinant DNA technology have also been used for some years in order to improve strains of the family 5 Enterobacteriaceae producing L-amino acids, by amplifying individual amino acid biosynthesis genes and investigating their effect on production.

Object of the Invention

The object of the invention is to provide new measures for 10 the improved enzymatic production of L-amino acids, in particular L-threonine.

Summary of the Invention

The present invention provides a process for the enzymatic production of L-amino acids, in particular L-threonine, 15 using microorganisms of the family Enterobacteriaceae that in particular already produce L-amino acids and in which the nucleotide sequence coding for the *dgsA* gene is attenuated.

Detailed Description of the Invention

20 Where L-amino acids or amino acids are mentioned hereinafter, this is understood to mean one or more amino acids including their salts, selected from the group comprising L-asparagine, L-threonine, L-serine, L-glutamate, L-glycine, L-alanine, L-cysteine, L-valine, L-25 methionine, L-isoleucine, L-leucine, L-tyrosine, L-phenylalanine, L-histidine, L-lysine, L-tryptophan and L-arginine. L-threonine is particularly preferred.

The term "attenuation" describes in this connection the reduction or switching off of the intracellular activity of 30 one or more enzymes (proteins) in a microorganism that are coded by the corresponding DNA, by using for example a weak

promoter or a gene or allele that codes for a corresponding enzyme with a low activity and/or that inactivates the corresponding enzyme (protein) or gene, and optionally combining these measures.

5 By means of these attenuation measures the activity or concentration of the corresponding protein is generally reduced to 0 to 75%, 0 to 50%, 0 to 25%, 0 to 10% or 0 to 5% of the activity or concentration of the wild type protein, or the activity or concentration of the protein in 10 the initial microorganism.

The process is characterized in that the following steps are carried-out:

- a) fermentation of microorganism of the family Enterobacteriaceae in which the dgsA gene is attenuated,
- 15 b) enrichment of the corresponding L-amino acid in the medium or in the cells of the microorganisms of the family Enterobacteriaceae, and
- c) isolation of the desired L-amino acid, in which 20 optionally constituents of the fermentation broth and/or the biomass in its entirety or parts thereof remain in the product.

The microorganisms that are the subject of the present invention can produce L-amino acids from glucose, sucrose, 25 lactose, fructose, maltose, molasses, optionally starch, optionally cellulose or from glycerol and ethanol. The microorganisms are members of the family Enterobacteriaceae selected from the genera Escherichia, Erwinia, Providencia and Serratia. The genera Escherichia and Serratia are 30 preferred. In the case of the genus Escherichia the species Escherichia coli may in particular be mentioned, and in the case of the genus Serratia the species Serratia marcescens may in particular be mentioned.

Suitable strains of the genus *Escherichia*, in particular those of the species *Escherichia coli*, that produce in particular L-threonine include for example:

- 5 *Escherichia coli* TF427
- Escherichia coli* H4578
- Escherichia coli* KY10935
- Escherichia coli* VNIIgenetika MG442
- Escherichia coli* VNIIgenetika M1
- Escherichia coli* VNIIgenetika 472T23
- 10 *Escherichia coli* BKIIM B-3996
- Escherichia coli* kat 13
- Escherichia coli* KCCM-10132

Suitable strains of the genus *Serratia*, in particular of the species *Serratia marcescens*, that produce L-threonine 15 include for example:

- Serratia marcescens* HNr21
- Serratia marcescens* TLr156
- Serratia marcescens* T2000

Strains of the family of *Enterobacteriaceae* producing L-threonine preferably have, *inter alia*, one or more of the 20 genetic or phenotype features selected from the following group: resistance to α -amino- β -hydroxyvaleric acid, resistance to thialysine, resistance to ethionine, resistance to α -methylserine, resistance to diaminosuccinic 25 acid, resistance to α -aminobutyric acid, resistance to borrelidin, resistance to rifampicin, resistance to valine analogues such as for example valine hydroxamate, resistance to purine analogues such as for example 6-dimethylaminopurine, need for L-methionine, optionally 30 partial and compensatable need for L-isoleucine, need for meso-diaminopimelic acid, auxotrophy with regard to threonine-containing dipeptides, resistance to L-threonine, resistance to L-homoserine, resistance to L-lysine, resistance to L-methionine, resistance to L-glutamic acid,

resistance to L-aspartate, resistance to L-leucine,
resistance to L-phenylalanine, resistance to L-serine,
resistance to L-cysteine, resistance to L-valine,
sensitivity to fluoropyruvate, defective threonine
5 dehydrogenase, optionally ability to utilise sucrose,
enhancement of the threonine operon, enhancement of
homoserine dehydrogenase, I-aspartate kinase I, preferably
of the feedback-resistant form, enhancement of homoserine
kinase, enhancement of threonine synthase, enhancement of
10 aspartate kinase, optionally of the feedback-resistant
form, enhancement of aspartate semialdehyde dehydrogenase,
enhancement of phosphoenol pyruvate carboxylase, optionally
of the feedback-resistant form, enhancement of phosphoenol
pyruvate synthase, enhancement of transhydrogenase,
15 enhancement of the RhtB gene product, enhancement of the
RhtC gene product, enhancement of the YfiK gene product,
enhancement of a pyruvate carboxylase, and attenuation of
acetic acid formation.

It has now been found that microorganisms of the family
20 Enterobacteriaceae after attenuation, in particular after
switching off the dgsA gene, produce L-amino acids, in
particular L-threonine, in an improved way.

The nucleotide sequences of the *Escherichia coli* genes
belong to the prior art and may also be obtained from the
25 genome sequence of *Escherichia coli* published by Blattner
et al. (Science 277, 1453 - 1462 (1997)).

The dgsA gene is described *inter alia* by the following
data:

Designation: Regulator of the phosphotransferase system
30 EC-No.: -
Reference: Hosono et al.; Bioscience, Biotechnology
and Biochemistry 59, 256-261 (1995) Morris
et al.; Journal of Bacteriology 163, 785-786
(1985)

Accession No.: AE000255

Note: The *dgsA* gene is also designated as *mlc* gene in the prior art.

Apart from the described *dgsA* gene, alleles of the gene may 5 be used that result from the degeneracy of the genetic code or from functionally neutral sense mutations, the activity of the protein not being substantially altered.

In order to achieve an attenuation the expression of the gene or the catalytic properties of the enzyme proteins may 10 for example be reduced or switched off. Optionally both measures may be combined.

The gene expression may be reduced by suitable culture conditions, by genetic alteration (mutation) of the signal structures of the gene expression, or also by antisense-RNA 15 techniques. Signal structures of the gene expression are for example repressor genes, activator genes, operators, promoters, attenuators, ribosome-binding sites, the start codon and terminators. The person skilled in the art may find relevant information in, *inter alia*, articles by 20 Jensen and Hammer (Biotechnology and Bioengineering 58: 191-195 (1998)), by Carrier and Keasling (Biotechnology Progress 15, 58-64 (1999)), Franch and Gerdes (Current Opinion in Microbiology 3, 159-164 (2000)) and in known textbooks of genetics and molecular biology, such as for 25 example the textbook by Knippers ("Molekulare Genetik", 6th Edition, Georg Thieme Verlag, Stuttgart, Germany, 1995) or that by Winnacker ("Gene and Klone", VCH Verlagsgesellschaft, Weinheim, Germany, 1990).

Mutations that lead to a change or reduction of the 30 catalytic properties of enzyme proteins are known from the prior art. As examples there may be mentioned the work by Qiu and Goodman (Journal of Biological Chemistry 272: 8611-8617 (1997)), Yano et al. (Proceedings of the National Academy of Sciences, USA 95, 5511-5515 (1998)), Wente and

Schachmann (Journal of Biological Chemistry 266, 20833-20839 (1991). Detailed information may be obtained from known textbooks on genetics and molecular biology, such as for example that by Hagemann ("Allgemeine Genetik", Gustav Fischer Verlag, Stuttgart, 1986).

Suitable mutations include transitions, transversions, insertions and deletions. Depending on the action of the amino acid exchange on the enzyme activity, one speaks of missense mutations or nonsense mutations. Insertions or 10 deletions of at least one base pair in a gene lead to frame shift mutations, which in turn lead to the incorporation of false amino acids or the premature termination of a translation: If as a result of the mutation a stop codon is formed in the coding region, this also leads to a 15 premature termination of the translation. Deletions of several codons typically lead to a complete disruption of the enzyme activity. Details regarding the production of such mutations belong to the prior art and may be obtained from known textbooks on genetics and molecular biology, 20 such as for example the textbook by Knippers ("Molekulare Genetik", 6th Edition, Georg Thieme Verlag, Stuttgart, Germany, 1995), that by Winnacker ("Gene und Klone", VCH Verlagsgesellschaft, Weinheim, Germany, 1990) or that by Hagemann ("Allgemeine Genetik", Gustav Fischer Verlag, 25 Stuttgart, 1986).

Suitable mutations in the genes such as for example deletion mutations may be incorporated by gene and/or allele exchange in suitable strains.

A conventional method is the method of gene exchange by 30 means of a conditionally replicating pSC101 derivate pMAK705 described by Hamilton et al. (Journal of Bacteriology 171, 4617 - 4622 (1989)). Other methods described in the prior art, such as for example that of Martinez-Morales et al. (Journal of Bacteriology 181, 7143-

7148 (1999)) or that of Boyd et al. (Journal of Bacteriology 182, 842-847 (2000)) may likewise be used.

It is also possible to transfer mutations in the respective genes or mutations relating to the expression of the 5 relevant genes, by conjugation or transduction into various strains.

Furthermore for the production of L-amino acids, in particular L-threonine, using strains of the family Enterobacteriaceae it may be advantageous in addition to 10 the attenuation of the dgsA gene also to enhance one or more enzymes of the known threonine biosynthesis pathway or enzymes of anaplerotic metabolism or enzymes for the production of reduced nicotinamide-adenine-dinucleotide phosphate.

15 The term "enhancement" describes in this connection the raising of the intracellular activity of one or more enzymes or proteins in a microorganism that are coded by the corresponding DNA, by for example increasing the number of copies of the gene or genes, using a strong promoter or 20 a gene that codes for a corresponding enzyme or protein having a high activity, and optionally by combining these measures.

By means of the aforementioned enhancement measures, in particular overexpression, the activity or concentration of 25 the corresponding protein is in general raised by at least 10%, 25%, 50%, 75%, 100%, 150%, 200%, 300%, 400% or 500%, at most up to 1000% or 2000% referred to that of the wild type protein and/or the activity or concentration of the protein in the initial microorganism.

30 Thus, one or more of the genes selected from the following group may for example by simultaneously enhanced, in particular overexpressed:

- the thrABC operon coding for aspartate kinase, homoserine dehydrogenase, homoserine kinase and threonine synthase (US-A-4,278,765),
- the pyc gene coding for pyruvate carboxylase 5 (DE-A-19 831 609),
- the pps gene coding for phosphoenol pyruvate synthase (Molecular and General Genetics 231:332 (1992)),
- the ppc gene coding for phosphoenol pyruvate carboxylase (Gene 31:279-283 (1984)),
- 10 • the genes pntA and pntB coding for transhydrogenase (European Journal of Biochemistry 158:647-653 (1986)),
- the gene rhtB imparting homoserine resistance (EP-A-0 994 190),
- 15 • the mqo gene coding for malate:quinone oxidoreductase (DE 100 348 33.5),
- the gene rhtC imparting threonine resistance (EP-A-1 013 765), and
- the thrE gene of *Corynebacterium glutamicum* coding for threonine export (DE 100 264 94.8).

20 The use of endogenous genes is in general preferred. The term "endogenous genes" or "endogenous nucleotide sequences" is understood to mean the genes or nucleotide sequences present in the population of a species.

Furthermore for the production of L-amino acids, in 25 particular L-threonine, it may be advantageous in addition to the attenuation of the dgsA gene also to attenuate, in particular to switch off or reduce the expression of one or more of the genes selected from the following group:

- the *tdh* gene coding for threonine dehydrogenase (Ravnikar and Somerville, *Journal of Bacteriology* 169, 4716-4721 (1987)),
- the *mdh* gene coding for malate dehydrogenase (E.C. 1.1.1.37) (Vogel et al., *Archives in Microbiology* 149, 36-42 (1987)),
- the gene product of the open reading frame (orf) *yjfa* (Accession Number AAC77180 of the National Center for Biotechnology Information (NCBI, Bethesda, MD, USA)),
- 10 • the gene product of the open reading frame (orf) *ytfP* (Accession Number AAC77179 des National Center for Biotechnology Information (NCBI, Bethesda, MD, USA)),
- the *pckA* gene coding for the enzyme phosphoenol pyruvate carboxykinase (Medina et al. (*Journal of Bacteriology* 172, 7151-7156 (1990))),
- 15 • the *poxB* gene coding for pyruvate oxidase (Grabau and Cronan (*Nucleic Acids Research* 14 (13), 5449-5460 (1986))),
- the *fruR* gene coding for the fructose repressor: 20 (Jahreis et al., *Molecular and General Genetics* 226, 332-336 (1991) and Accession No.: AE000118), and
- the *aceA* gene for isocitrate lyase (EC-No.: 4.1.3.1) kodierende (Matsuoko and McFadden; *Journal of Bacteriology* 170, 4528-4536 (1988) and Accession No.: 25 AE000474)

Furthermore for the production of L-amino acids, in particular L-threonine, it may be advantageous in addition to the attenuation of the *dgsA* gene also to switch off undesirable secondary reactions (Nakayama: "Breeding of 30 Amino Acid Producing Microorganisms", in: *Overproduction of*

Microbial Products, Krumphanzl, Sickyta, Vanek (eds.), Academic Press, London, UK, 1982).

The microorganisms produced according to the invention may be cultivated in a batch process (batch cultivation), in a 5 fed batch process (feed process) or in a repeated fed batch process (repetitive feed process). A summary of known cultivation methods is described in the textbook by Chmiel (Bioprozesstechnik 1. Einführung in die Bioverfahrenstechnik (Gustav Fischer Verlag, Stuttgart, 10 1991)) or in the textbook by Storhas (Bioreaktoren and periphere Einrichtungen (Vieweg Verlag, Brunswick/Wiesbaden, 1994)).

The culture medium to be used must appropriately satisfy the requirements of the respective strains. Descriptions 15 of culture media of various microorganisms are contained in the handbook "Manual of Methods for General Bacteriology" of the American Society for Bacteriology (Washington D.C., USA, 1981).

As carbon sources, sugars and carbohydrates such as for 20 example glucose, sucrose, lactose, fructose, maltose, molasses, starch and optionally cellulose, oils and fats such as for example soya bean oil, sunflower oil, groundnut oil and coconut oil, fatty acids such as for example palmitic acid, stearic acid and linoleic acid, alcohols 25 such as for example glycerol and ethanol, and organic acids such as for example acetic acid, may be used. These substances may be used individually or as a mixture.

As nitrogen source, organic nitrogen-containing compounds such as peptones, yeast extract, meat extract, malt 30 extract, maize starch water, soya bean flour and urea or inorganic compounds such as ammonium sulfate, ammonium chloride, ammonium phosphate, ammonium carbonate and ammonium nitrate may be used. The nitrogen sources may be used individually or as a mixture.

As phosphorus source, phosphoric acid, potassium dihydrogen phosphate or dipotassium hydrogen phosphate or the corresponding sodium-containing salts may be used. The culture medium must furthermore contain salts of metals, 5 such as for example magnesium sulfate or iron sulfate, that are necessary for growth. Finally, essential growth promoters such as amino acids and vitamins may be used in addition to the aforementioned substances. Apart from these, suitable precursors may be added to the culture 10 medium. The aforementioned starting substances may be added to the culture in the form of a single batch or may be metered in in an appropriate manner during the cultivation.

In order to regulate the pH of the culture basic compounds 15 such as sodium hydroxide, potassium hydroxide, ammonia or ammonia water, or acidic compounds such as phosphoric acid or sulfuric acid are used as appropriate. In order to control foam formation antifoaming agents such as for example fatty acid polyglycol esters may be used. In order 20 to maintain the stability of plasmids, suitable selectively acting substances, for example antibiotics, may be added to the medium. In order to maintain aerobic conditions, oxygen or oxygen-containing gas mixtures such as for example air are fed into the culture. The temperature of 25 the culture is normally 25°C to 45°C, and preferably 30°C to 40°C. Cultivation is continued until a maximum amount of L-amino acids (or L-threonine) has been formed. This target is normally achieved within 10 hours to 160 hours.

The L-amino acids may be analyzed by anion exchange 30 chromatography followed by ninhydrin derivation, as described by Spackman et al. (Analytical Chemistry, 30, (1958), 1190), or by reversed phase HPLC, as described by Lindroth et al. (Analytical Chemistry (1979) 51: 1167-1174).

The process according to the invention can be used for the enzymatic production of L-amino acids, such as for example L-threonine, L-isoleucine, L-valine, L-methionine, L-homoserine and L-lysine, in particular L-threonine.

5 A pure culture of the *Escherichia coli* K-12 strain DH5 α /pMAK705 was filed as DSM 13720 on 08 September 2000 at the German Collection for Microorganisms and Cell Cultures (DSMZ, Brunswick, Germany) according to the Budapest Convention.

10 The present invention is described in more detail hereinafter with the aid of examples of implementation.

The isolation of plasmid DNA from *Escherichia coli* as well as all techniques for the restriction, Klenow treatment and alkaline phosphatase treatment are carried out according to 15 Sambrook et al. (*Molecular Cloning - A Laboratory Manual* (1989) Cold Spring Harbor Laboratory Press). The transformation of *Escherichia coli* is, unless otherwise described, carried out according to Chung et al. (*Proceedings of the National Academy of Sciences of the 20 United States of America, USA* (1989) 86: 2172-2175).

The incubation temperature in the production of strains and transformants is 37°C. In the gene exchange process according to Hamilton et al., temperatures of 30°C and 44°C are used.

25 Example 1

Construction of the deletion mutation of the *dgsA* gene

Parts of the gene regions lying upstream and downstream of the *dgsA* gene and parts of the 5'-region and 3'-region of the *dgsA* gene are amplified from *Escherichia coli* K12 using 30 the polymerase chain reaction (PCR) as well as synthetic oligonucleotides. Starting from the nucleotide sequence of the *dgsA* gene and sequences in *E. coli* K12 MG1655 (SEQ ID

No. 1, Accession Number AE000255) lying upstream and downstream, the following PCR primers are synthesised (MWG Biotech, Ebersberg, Germany):

dgsA'5'-1: 5' - CGAATGTAACGCTGGCTGAA - 3' (SEQ ID No. 3)

5 dgsA'5'-2: 5' - TCCAGCAATGGCAAGTCATC - 3' (SEQ ID No. 4)

dgsA'3'-1: 5' - CAGCACATCAGCGTTGAGAG - 3' (SEQ ID No. 5)

dgsA'3'-2: 5' - GATCGCCTGAGCTGTTAGCA - 3' (SEQ ID No. 6)

The chromosomal *E. coli* K12 MG1655 DNA used for the PCR is isolated according to the manufacturer's instructions using 10 "Qiagen Genomic-tips 100/G" (QIAGEN, Hilden, Germany). A ca. 850 bp large DNA fragment from the 5'-region of the dgsA gene region (designated dgsA1) and a ca. 700 bp large DNA fragment from the 3'-region of the dgsA gene region (designated dgsA2) can be amplified with the specific 15 primers under standard PCR conditions (Innis et al. (1990) PCR Protocols. A Guide to Methods and Applications, Academic Press) with taq-DNA-polymerase (Gibco-BRL, Eggenstein, Germany). The PCR products are ligated according to the manufacturer's instructions in each case 20 with the vector pCR2.1TOPO (TOPO TA Cloning Kit, Invitrogen, Groningen, Netherlands) and transformed in the *E. coli* strain TOP10F'. The selection of plasmid-carrying cells is carried out on LB agar to which 50 µg/ml of ampicillin has been added. After the plasmid DNA isolation 25 the vector pCR2.1TOPOdgsA2 is cleaved with the restriction enzymes EcoRI and XbaI, and the dgsA2 fragment after separation in 0.8% agarose gel is isolated using the QIAquick Gel Extraction Kit (QIAGEN, Hilden, Germany). After the plasmid DNA isolation the vector pCR2.1TOPOdgsA1 30 is cleaved with the enzymes EcoRV and XbaI and ligated with the isolated dgsA2 fragment. The *E. coli* strain DH5α is transformed with the ligation batch and plasmid-carrying cells are selected on LB agar to which 50 µg/ml of

ampicillin has been added. After the plasmid DNA isolation, those plasmids in which the mutagenic DNA sequence shown in SEQ ID No. 7 is present in cloned form are detected by control cleavage with the enzymes HindIII and XbaI. One of the plasmids is designated 5 pCR2.1TOPOΔdgsA.

Example 2

Construction of the exchange vector pMAK705ΔdgsA

The dgsA allele described in Example 1 is isolated from the 10 vector pCR2.1TOPOΔdgsA after restriction with the enzymes HindIII and XbaI and separation in 0.8% agarose gel, and is ligated with the plasmid pMAK705 (Hamilton et al. (1989) Journal of Bacteriology 171, 4617 - 4622), that had been digested with the enzymes HindIII and XbaI. The ligation 15 batch is transformed in DH5 α and plasmid-carrying cells are selected on LB agar to which 20 μ g/ml of chloramphenicol have been added. The successful cloning is detected after plasmid DNA isolation and cleavage with the enzymes HindIII and XbaI. The resultant exchange vector pMAK705ΔdgsA (= 20 pMAK705deltadgsA) is shown in Fig. 1.

Example 3

Site-specific mutagenesis of the dgsA gene in the E. coli strain MG442

The E. coli strain MG442 producing L-threonine is described 25 in patent specification US-A- 4,278,765 and is filed as CMIM B-1628 at the Russian National Collection for Industrial Microorganisms (VKPM, Moscow, Russia).

For the exchange of the chromosomal dgsA gene by the plasmid-coded deletion construct, MG442 is transformed 30 with the plasmid pMAK705ΔdgsA. The gene exchange is carried out by the selection process described by Hamilton et al. (1989) Journal of Bacteriology 171, 4617 - 4622) and

is verified by standard PCR methods (Innis et al. (1990) PCR Protocols. A guide to methods and applications, Academic Press) with the following oligonucleotide primers:

dgsA'5'-1: 5' - CGAATGTAACGCTGGCTGAA - 3' (SEQ ID No. 3)

5 dgsA'3'-2: 5' - GATCGCCTGAGCTGTTAGCA - 3' (SEQ ID No. 6)

After the exchange the form of the Δ dgsA allele shown in SEQ ID No. 8 is present in MG442. The strain obtained is designated MG442 Δ dgsA.

Example 4

10 Production of L-threonine using the strain MG442 Δ dgsA

MG442 Δ dgsA is cultivated on minimal medium having the following composition: 3.5 g/l $\text{Na}_2\text{HPO}_4 \cdot 2\text{H}_2\text{O}$, 1.5 g/l KH_2PO_4 , 1 g/l NH_4Cl , 0.1 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 2 g/l glucose and 20 g/l agar. The formation of L-threonine is checked in batch 15 cultures of 10 ml that are contained in 100 ml Erlenmeyer flasks. For this, 10 ml of preculture medium of the following composition: 2 g/l yeast extract, 10 g/l $(\text{NH}_4)_2\text{SO}_4$, 1 g/l KH_2PO_4 , 0.5 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 15 g/l CaCO_3 , 20 g/l glucose are inoculated and incubated for 16 hours at 20 37°C and 180 rpm in an ESR incubator from Kühner AG (Birsfelden, Switzerland). 250 μl of this preculture are reinoculated in 10 ml of production medium (25 g/l $(\text{NH}_4)_2\text{SO}_4$, 2 g/l KH_2PO_4 , 1 g/l $\text{MgSO}_4 \cdot 7\text{H}_2\text{O}$, 0.03 g/l $\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$, 0.018 g/l $\text{MnSO}_4 \cdot 1\text{H}_2\text{O}$, 30 g/l CaCO_3 and 20 g/l glucose) and 25 incubated for 48 hours at 37°C. After incubation the optical density (OD) of the culture suspension is measured with an LP2W photometer from the Dr. Lange company (Dusseldorf, Germany) at a measurement wavelength of 660 nm.

30 The concentration of formed L-threonine is then determined in the sterile-filtered culture supernatant using an amino acid analyzer from Eppendorf-BioTronik (Hamburg, Germany)

by ion exchange chromatography and post-column reaction with ninhydrin detection.

The result of the test is given in Table 1.

Table 1

Strain	OD (660 nm)	L- threonine
MG442	6.0	1.5
MG442ΔdgsA	6.5	1.8

5

Brief Description of the Figure:

- Fig. 1: pMAK705ΔdgsA (= pMAK705deltadgsA)

Length data are given as approximate values. The abbreviations and acronyms used have the following
10 meanings:

- cat: chloramphenicol resistance gene
- rep-ts: temperature-sensitive replication region of the plasmid pSC101
- dgsA1: part of the 5'-region of the dgsA gene and of
15 the upstream-lying region
- dgsA2: part of the 3'-region of the dgsA gene and of the downstream-lying region

The abbreviations for the restriction enzymes have the following meanings:

20 • BamHI: restriction endonuclease from *Bacillus amyloliquefaciens*

- BglIII: restriction endonuclease from *Bacillus globigii*
- ClaI: restriction endonuclease from *Caryphanon latum*
- EcoRI: restriction endonuclease from *Escherichia coli*
- 5 • EcoRV: restriction endonuclease from *Escherichia coli*
- HindIII: restriction endonuclease from *Haemophilus influenzae*
- KpnI: restriction endonuclease from *Klebsiella pneumoniae*
- 10 • PstI: restriction endonuclease from *Providencia stuartii*
- PvuI: restriction endonuclease from *Proteus vulgaris*
- SacI: restriction endonuclease from *Streptomyces achromogenes*
- 15 • SalI: restriction endonuclease from *Streptomyces albus*
- SmaI: restriction endonuclease from *Serratia marcescens*
- 20 • SphI: restriction endonuclease from *Streptomyces phaeochromogenes*
- SspI: restriction endonuclease from *Sphaerotilus species*
- XbaI: restriction endonuclease from *Xanthomonas badrii*
- 25 • XhoI: restriction endonuclease from *Xanthomonas holcicola*

What is Claimed is:

1. Process for the production of L-amino acids, in particular L-threonine, wherin the following steps are carried out:
 - 5 a) fermentation of the microorganisms of the family Enterobacteriaceae producing the desired L-amino acid, in which the dgsA gene or nucleotide sequences coding therefor are attenuated, in particular are switched off,
 - 10 b) enrichment of the L-amino acid in the medium or in the cells of the microorganisms, and
 - 15 c) isolation of the L-amino acid, in which optionally constituents of the fermentation broth and/or the biomass in its entirety or portions thereof remain in the product.
2. Process according to claim 1, wherein microorganisms are used in which in addition further genes of the biosynthesis pathway of the desired L-amino acid are enhanced.
- 20 3. Process according to claim 1, wherein microorganisms are used in which the metabolic pathways that reduce the formation of the desired L-amino acid are at least partially switched off.
4. Process according to claim 1, wherein the expression of 25 the polynucleotide(s) that codes/code for the dgsA gene is attenuated, in particular is switched off.
5. Process according to claim 1, wherein the regulatory and/or catalytic properties of the polypeptide (enzyme protein) for which the polynucleotide dgsA codes are 30 reduced.

6. Process according to claim 1, wherein for the production of L-amino acids microorganisms of the family Enterobacteriaceae are fermented in which at the same time one or more of the genes selected from the following group is enhanced, in particular overexpressed:
 - 6.1 the thrABC operon coding for aspartate kinase, homoserine dehydrogenase, homoserine kinase and threonine synthase,
 - 10 6.2 the pyc gene coding for pyruvate carboxylase,
 - 6.3 the pps gene coding for phosphoenol pyruvate synthase,
 - 6.4 the ppc gene coding for phosphoenol pyruvate carboxylase,
 - 15 6.5 the genes pntA and pntB coding for transhydrogenase,
 - 6.6 the gene rhtB imparting homoserine resistance,
 - 6.7 the mqo gene coding for malate:quinone oxidoreductase,
 - 20 6.8 the gene rhtC imparting threonine resistance, and
 - 6.9 the thrE gene coding for threonine export.
7. Process according to claim 1, wherein for the production of L-amino acids microorganisms of the family Enterobacteriaceae are fermented in which at the same time one or more of the genes selected from the following group is attenuated, in particular switched off, or the expression is reduced:
 - 7.1 the tdh gene coding for threonine dehydrogenase,
 - 7.2 the mth gene coding for malate dehydrogenase,

7.3 the gene product of the open reading frame (orf) yjfA,

7.4 the gene product of the open reading frame (orf) ytfP,

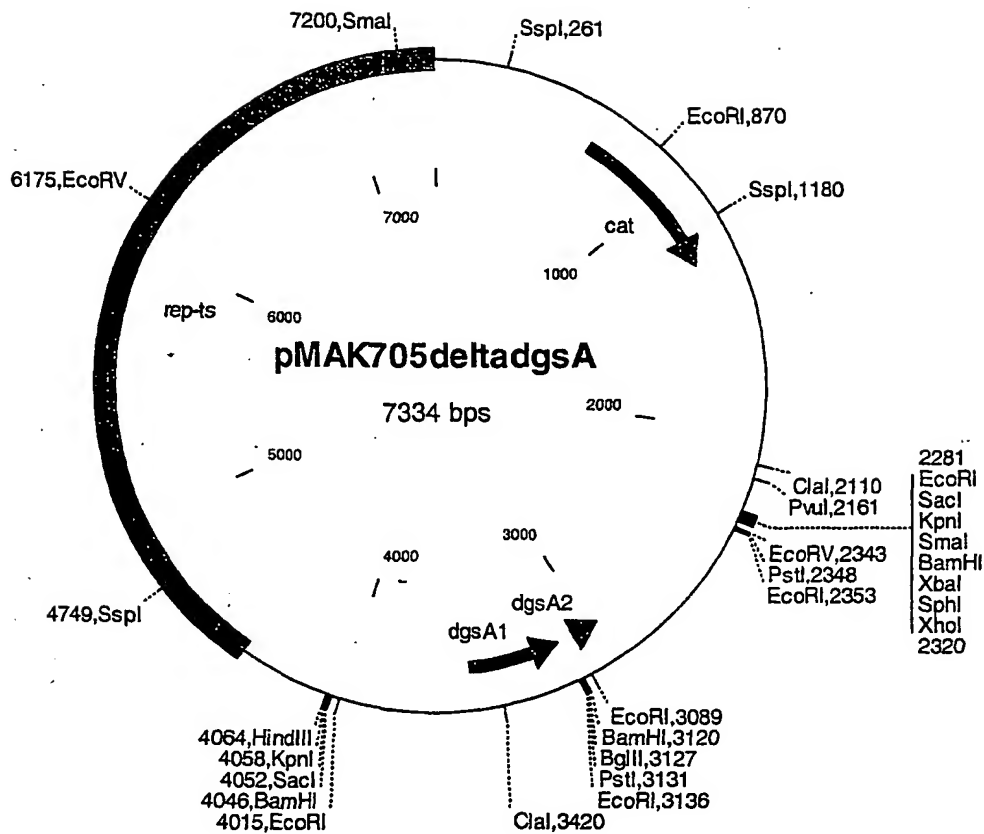
5 7.5 the pckA gene coding for phosphoenol pyruvate carboxykinase,

7.6 the poxB gene coding for pyruvate oxidase,

7.7 the fruR gene coding for the fructose repressor,

7.8 the aceA gene coding for isocitrate lyase.

Fig. 1:



SEQUENCE LISTING

5 <110> Degussa AG
 10 <120> Process for the production of L-amino acids using strains of the
 family Enterobacteriaceae containing an attenuated dgsA-Gen
 15 <130> 020003 BT
 20 <160> 8
 25 <170> PatentIn version 3.1
 30 <210> 1
 <211> 2306
 <212> DNA
 <213> Escherichia coli
 35 <220>
 <221> CDS
 <222> (485)..(1705)
 <223> dgsA gene
 40 <400> 1
 cgaatgtaac gctggctgaa ctggcgaaag aaccctttgt ctttttgat cccgacgtcg 60
 ggacagggct gtatgacgat attctcgggc tcatgcgacg ttaccatttg acgcccgtca 120
 45 tcactcagga ggtgggcgag gcaatgacca tcatcggtct ggttccgccc ggtctgggtg 180
 tttcaatttt gcctgcgtca tttaaacgtg ttcaagctcaa cgaaatgcgc tgggtgcccga 240
 50 ttgctgaaga ggatgcggtt tctgaaatgt ggttggtctg gccgaaacat catgaacaaa 300
 gtccggctgc gcgttaacttt cgtattcatc tgctgaatgc tctcaggtga gggaaatttc 360
 55 agcgaaaaag cccgaaaaat gtgctgttaa tcacatgcct aagtaaaaat ttgacgacac 420
 60 gtattgaagt gtttcaccat agcctacaga ttatccgga gcgaaaaat atagggagta 480
 tgcg gtg gtt gct gaa aac cag cct ggg cac att gat caa ata aag cag 529
 Met Val Ala Glu Asn Gln Pro Gly His Ile Asp Gln Ile Lys Gln
 1 5 10 15
 65 acc aac gcg ggc gcg gtt tat cgc ctg att gat cag ctt ggt cca gtc 577
 Thr Asn Ala Gly Ala Val Tyr Arg Leu Ile Asp Gln Leu Gly Pro Val
 20 25 30
 70 tcg cgt atc gat ctt tcc cgt ctg gcg caa ctg gct cct gcc agt atc 625
 Ser Arg Ile Asp Leu Ser Arg Leu Ala Gln Leu Ala Pro Ala Ser Ile
 35 40 45
 75 act aaa att gtc cgt gag atg ctc gaa gca cac ctg gtg caa gag ctg 673
 Thr Lys Ile Val Arg Glu, Met Leu Glu Ala His Leu Val Gln Glu Leu
 50 55 60
 80 gaa atc aaa gaa gcg ggg aac cgt ggc cgt ccg gcg gtg ggg ctg gtg 721
 Glu Ile Lys Glu Ala Gly Asn Arg Gly Arg Pro Ala Val Gly Leu Val
 65 70 75
 85 gtt gaa act gaa gcc tgg cac tat ctt tct ctg cgc att agt cgc ggg 769
 Val Glu Thr Glu Ala Trp His Tyr Leu Ser Leu Arg Ile Ser Arg Gly
 80 85 90 95

65

	gag att ttc ctt gct ctg cgc gat ctg agc agc aaa ctg gtg gtg gaa Glu Ile Phe Leu Ala Leu Arg Asp Leu Ser Ser Lys Leu Val Val Glu 100 105 110	817
5	gag tcg cag gaa ctg gcg tta aaa gat gac ttg cca ttg ctg gat cgt Glu Ser Gln Glu Leu Ala Leu Lys Asp Asp Leu Pro Leu Leu Asp Arg 115 120 125	865
10	att att tcc cat atc gat cag ttt ttt atc cgc cac cag aaa aaa ctt Ile Ile Ser His Ile Asp Gln Phe Ile Arg His Gln Lys Lys Leu 130 135 140	913
15	gag cgt cta act tcg att gcc ata acc ttg ccg gga att att gat acg Glu Arg Leu Thr Ser Ile Ala Ile Thr Leu Pro Gly Ile Ile Asp Thr 145 150 155	961
20	gaa aat ggt att gta cat cgc atg ccg ttc tac gag gat gta aaa gag Glu Asn Gly Ile Val His Arg Met Pro Phe Tyr Glu Asp Val Lys Glu 160 165 170 175	1009
25	atg ccg ctc ggc gag gcg ctg gag cag cat acc ggc gtt ccg gtt tat Met Pro Leu Gly Glu Ala Leu Glu Gln His Thr Gly Val Pro Val Tyr 180 185 190	1057
30	att cag cat gat atc agc gca tgg acg atg gca gag gcc ttg ttt ggt Ile Gln His Asp Ile Ser Ala Trp Thr Met Ala Glu Ala Leu Phe Gly 195 200 205	1105
35	gcc tca cgc ggg gcg cgc gat gtg att cag gtg gtt atc gat cac aac Ala Ser Arg Gly Ala Arg Asp Val Ile Gln Val Val Ile Asp His Asn 210 215 220	1153
40	gtg ggg gcg ggc gtc att acc gat ggt cat ctg cta cac gca ggc agc Val Gly Ala Gly Val Ile Thr Asp Gly His Leu Leu His Ala Gly Ser 225 230 235	1201
45	agt agt ctc gtg gaa ata ggc cac aca cag gtc gac ccg tat ggg aaa Ser Ser Leu Val Glu Ile Gly His Thr Gln Val Asp Pro Tyr Gly Lys 240 245 250 255	1249
50	cgc tgt tat tgc ggg aat cac ggc tgc ctc gaa acc atc gcc agc gtg Arg Cys Tyr Cys Gly Asn His Gly Cys Leu Glu Thr Ile Ala Ser Val 260 265 270	1297
55	gac agt att ctt gag ctg gca cag ctg cgt ctt aat caa tcc atg agc Asp Ser Ile Leu Glu Leu Ala Gln Leu Arg Leu Asn Gln Ser Met Ser 275 280 285	1345
60	tcg atg tta cat gga caa ccg tta acc gtg gac tca ttg tgt cag gcg Ser Met Leu His Gly Gln Pro Leu Thr Val Asp Ser Leu Cys Gln Ala 290 295 300	1393
65	gca ttg cgc ggc gat cta ctg gca aaa gac atc att acc ggg gtg ggc Ala Leu Arg Gly Asp Leu Leu Ala Lys Asp Ile Ile Thr Gly Val Gly 305 310 315	1441
70	gcg cat gtc ggg cgc att ctt gcc atc atg gtg aat tta ttt aac cca Ala His Val Gly Arg Ile Leu Ala Ile Met Val Asn Leu Phe Asn Pro 320 325 330 335	1489
75	caa aaa ata ctg att ggc tca ccg tta agt aaa gcg gca gat atc ctc Gln Lys Ile Leu Ile Gly Ser Pro Leu Ser Lys Ala Ala Asp Ile Leu 340 345 350	1537

	ttc ccg gtc atc tca gac agc atc cgt cag cag gcc ctt cct gcg tat	1585
	Phe Pro Val Ile Ser Asp Ser Ile Arg Gln Gln Ala Leu Pro Ala Tyr	
	355 360 365	
5	agt cag cac atc agc gtt gag agt act cag ttt tct aac cag ggc acg	1633
	Ser Gln His Ile Ser Val Glu Ser Thr Gln Phe Ser Asn Gln Gly Thr	
	370 375 380	
10	atg gca ggc gct gca ctg gta aaa gac gcg atg tat aac ggt tct ttg	1681
	Met Ala Gly Ala Ala Leu Val Lys Asp Ala Met Tyr Asn Gly Ser Leu	
	385 390 395	
15	ttg att cgt ctg ttg cag ggt taa catttttaa ctgttctacc aaaatttgcg	1735
	Leu Ile Arg Leu Leu Gln Gly	
	400 405	
	ctatctcaat ttggccagg aaagcataac ttagacttc aaggttaatt attttcctgg	1795
20	tttatatttgc tgaagcataa cggggagtt agtgatgctg aagcgttct ttattaccgg	1855
	tacagacact tctgttaggaa aaacgggtggt ttcccgcgca ttgctacaag cgtagcctc	1915
	ccagggaaaa acgggtgcgg gatataaacc cgtagcgaag gggagcaaag agacacccga	1975
25	agggctgcgt aataaaagatg ccctggtggtt gcagagtgtt tcaaccatcg aactgcctta	2035
	tgaaggcagtt aatcctatcg cgtaagcga agaagaaagt agcgtggcgc acagttgccc	2095
30	aatcaattac accctcattt caaacggcct ggcaaaccctg accgaaaaag tcgatcatgt	2155
	cgtggtagaa gggactggcg gctggcgag tctgatgaat gatttgcgtc cactctctga	2215
	atgggttagtgcaggaacaac tgccggtggtt gatggttgtc ggtattcagg aaggttgcatt	2275
35	taaccatgca ctgctaacag ctcaggcgat c	2306
	<210> 2	
	<211> 406	
40	<212> PRT	
	<213> Escherichia coli	
	<400> 2	
	Met Ala Glu Asn Gln Pro Gly His Ile Asp Gln Ile Lys Gln Thr	
45	1 5 10 15	
	Asn Ala Gly Ala Val Tyr Arg Leu Ile Asp Gln Leu Gly Pro Val Ser	
	20 25 30	
50	Arg Ile Asp Leu Ser Arg Leu Ala Gln Leu Ala Pro Ala Ser Ile Thr	
	35 40 45	
	Lys Ile Val Arg Glu Met Leu Glu Ala His Leu Val Gln Glu Leu Glu	
55	50 55 60	
	Ile Lys Glu Ala Gly Asn Arg Gly Arg Pro Ala Val Gly Leu Val Val	
	65 70 75 80	
60	Glu Thr Glu Ala Trp His Tyr Leu Ser Leu Arg Ile Ser Arg Gly Glu	
	85 90 95	
	Ile Phe Leu Ala Leu Arg Asp Leu Ser Ser Lys Leu Val Val Glu Glu	
	100 105 110	
65	Ser Gln Glu Leu Ala Leu Lys Asp Asp Leu Pro Leu Leu Asp Arg Ile	

	115	120	125
	Ile Ser His Ile Asp Gln Phe Phe Ile Arg His Gln Lys Lys Leu Glu		
	130	135	140
5	Arg Leu Thr Ser Ile Ala Ile Thr Leu Pro Gly Ile Ile Asp Thr Glu		
	145	150	155
	160		
10	Asn Gly Ile Val His Arg Met Pro Phe Tyr Glu Asp Val Lys Glu Met		
	165	170	175
	Pro Leu Gly Glu Ala Leu Glu Gln His Thr Gly Val Pro Val Tyr Ile		
	180	185	190
15	195 Gln His Asp Ile Ser Ala Trp Thr Met Ala Glu Ala Leu Phe Gly Ala		
	200	205	
	Ser Arg Gly Ala Arg Asp Val Ile Gln Val Val Ile Asp His Asn Val		
	210	215	220
20	Gly Ala Gly Val Ile Thr Asp Gly His Leu Leu His Ala Gly Ser Ser		
	225	230	235
	240		
25	Ser Leu Val Glu Ile Gly His Thr Gln Val Asp Pro Tyr Gly Lys Arg		
	245	250	255
	Cys Tyr Cys Gly Asn His Gly Cys Leu Glu Thr Ile Ala Ser Val Asp		
	260	265	270
30	Ser Ile Leu Glu Leu Ala Gln Leu Arg Leu Asn Gln Ser Met Ser Ser		
	275	280	285
	Met Leu His Gly Gln Pro Leu Thr Val Asp Ser Leu Cys Gln Ala Ala		
	290	295	300
35	Leu Arg Gly Asp Leu Leu Ala Lys Asp Ile Ile Thr Gly Val Gly Ala		
	305	310	315
	320		
40	His Val Gly Arg Ile Leu Ala Ile Met Val Asn Leu Phe Asn Pro Gln		
	325	330	335
	Lys Ile Leu Ile Gly Ser Pro Leu Ser Lys Ala Ala Asp Ile Leu Phe		
	340	345	350
45	355 Pro Val Ile Ser Asp Ser Ile Arg Gln Gln Ala Leu Pro Ala Tyr Ser		
	360	365	
	Gln His Ile Ser Val Glu Ser Thr Gln Phe Ser Asn Gln Gly Thr Met		
	370	375	380
50	Ala Gly Ala Ala Leu Val Lys Asp Ala Met Tyr Asn Gly Ser Leu Leu		
	385	390	395
	400		
55	Ile Arg Leu Leu Gln Gly		
	405		
	<210> 3		
	<211> 20		
	<212> DNA		
60	<213> Artificial sequence		
	<220>		
	<221> Primer		
	<222> (1)..(20)		
65	<223> dgsA '5`-1		

```

<400> 3
cgaatgtaac gctggctgaa
5
<210> 4
<211> 20
<212> DNA
<213> Artificial sequence
10
<220>
<221> Primer
<222> (1)..(20)
<223> dgsA '5'-2
15
<400> 4
tccagcaatg gcaagtcatc
20
<210> 5
<211> 20
<212> DNA
<213> Artificial sequence
25
<220>
<221> Primer
<222> (1)..(20)
<223> dgsA '3'-1
30
<400> 5
cagcacatca gcgttgagag
35
<210> 6
<211> 20
<212> DNA
<213> Artificial sequence
40
<220>
<221> Primer
<222> (1)..(20)
<223> dgsA '3'-2
45
<400> 6
gatcgccctga gctgttagca
50
<210> 7
<211> 1756
<212> DNA
<213> Escherichia coli
55
<220>
<221> misc_feature
<222> (1)..(1756)
<223>
60
<220>
<221> misc_feature
<222> (1)..(60)
<223> Technical DNA/ remainder polylinker sequence
65
<220>
<221> misc_feature
<222> (61)..(921)
<223> Part of the upstream-lying region and part of the 5'-region
      of the dgsA gene

```

```

<220>
<221> misc_feature
<222> (922)..(986)
<223> Technical DNA/ remainder polylinker sequence
5
<220>
<221> misc_feature
<222> (987)..(1704)
<223> Part of the 3'-region of the dgsA gene and part of the downstream-
10 lying region

<220>
<221> misc_feature
<222> (1705)..(1756)
15 <223> Technical DNA/ remainder polylinker sequence

<400> 7
    agcttggtag ctagtcggta tccactatgt aacggccggca gtgtgctggaa attcgccctt 60
20    cgaatgttaac gctggctgaa ctggcgaaag aaccctttgt ctttttgtat ccgcacgtcg 120
        ggacagggct gtatgacgat attctcggtc ttagtgcacg ttaccatttg acgcccgtca 180
25    tcactcagga ggtggcgag gcaatgacca tcatcggtct ggtttccggcc ggtctgggtg 240
        ttcaatttt gcctgcgtca. tttaaacgtt ttcagctaa cggaaatgcgc tgggtgccga 300
        ttgctgaaga ggatgcggtt tctgaaatgt ggttggcttg gccgaaacat catgaacaaaa 360
30    gtcgggctgc gcgttaacttt cgtattcatc tgctgaatgc tctcagggtga gggaaatttc 420
        agcgaaaaag cccgaaaaat gtgctgttaa tcacatgcct aagtaaaaaat ttgacgacac 480
35    gtattgaagt gcttcaccat agcctacaga ttatttcggaa ggcgaaaaat atagggagta 540
        tgccgggtt gctgaaaacc agcctggca cattgatcaa ataaagcaga ccaacgcggg 600
        cgcggtttat cgcctgattt atcagcttgg tccagtcgtcg cgtatcgatc tttccgtct 660
40    ggcgcaactg gctcctgcca gtatcactaa aattgtccgt gagatgcgtc aagcacacct 720
        ggtgcaagag ctggaaatca aagaagcggg gaaccgtggc cgtccggcgg tggggctgg 780
45    ggttggaaact gaagcctggc actatcttc tctgcgcatt agtgcgggg agatttcct 840
        tgctctgcgc gatctgagca gcaaaactggt ggtgaaagag tcgcaggaac tggcgtaaaa 900
        agatgacttg ccattgctgg aaagggcgaa ttctgcagat ctcggatcca cttagtaacgg 960
50    cccgcgtgt gctgaaattc gcccttcgc acatcagcgt tgagagtact cagttttctta 1020
        accagggcac gatggcaggc gctgcactgg taaaagacgc gatgtataac ggttctttgt 1080
55    tgattcgctc gttgcagggt taacatttt taactgttct accaaaattt ggcgtatctc 1140
        aatttgggcc agggaaagcat aacttagact ttcaagggtt attattttcc tggtttatat 1200
        ttgtgaagca taacgggtt gtttagtgcgt ctgaagcggt tctttattac cggtagac 1260
60    acttctgttag ggaaaacgggt ggttccgc gcattgcgtac aagcggttagc ctcccgaggaa 1320
        aaaacgggtt cgggatataa acccgtagcg aaggggagca aagagacacc cgaaggggctg 1380
        cgtataaaag atgcctgggt gttgcagagt gttcaaccca tcgaactgcc ttatgaagca 1440
65

```

gttaatccta tcgcgttaag cgaagaagaa agtagcgtgg cgcacagttg cccaatcaat 1500
tacaccctca tttcaaacgg cctggcaaac ctgaccgaaa aagtgcgtca tgcgtggta 1560
5 gaaggggactg gcggctggcg cagtctgatg aatgattgc gtccactctc tgaatgggta 1620
gtgcaggaac aactgccgtt gttgatggtt gtcggattc aggaagggtt cattaaccat 1680
gcactgctaa cagctcaggc gatcaaggc gaattctgca gatatccatc acactggcgg 1740
10 ccgctcgagc atgcat 1756

<210> 8
<211> 559
15 <212> DNA
<213> Escherichia coli

<220>
<221> misc_feature
20 <222> (1)..(559)
<223>

<220>
<221> misc_feature
25 <222> (1)..(3)
<223> Start codon of the delta dgsA allele

<220>
<221> misc_feature
30 <222> (1)..(377)
<223> 5'-region of the delta dgsA allele

<220>
<221> misc_feature
35 <222> (378)..(442)
<223> Technical DNA/ remainder polylinker sequence

<220>
<221> misc_feature
40 <222> (443)..(556)
<223> 3'-region of the delta dgsA allele

<220>
<221> misc_feature
45 <222> (557)..(559)
<223> Stop codon of the delta dgsA allele

<400> 8
50 gtggttgctg aaaaccagcc tggcacatt gatcaaataa agcagaccaa cgccggcg 60
gtttatcgcc tgattgatca gcttggtcca gtctcgctgatcgatcttc ccgtctggcg 120
caactggctc ctgccagtat cactaaaatt gtccgtgaga tgctcgaagc acacctggtg 180
55 caagagctgg aaatcaaaga agcggggAAC cgtggccgtc cggcggtggg gctgggtgg 240
gaaaactgaag cctggcacta tctttctctg cgcattagtc gcggggagat tttcccttgct 300
60 ctgcgcgatc tgagcagcaa actgggtggta gaagagtgc aggaactggc gttaaaaagat 360
gacttgccat tgctggaaag ggcgaattct gcagatctg gatccactag taacggccgc 420
cagtgtgctg gaattcgccc ttcagcacat cagcgtttagt agtactcagt tttctaaacca 480
65 gggcacgatg gcaggcgctg cactggtaaa agacgcgtatg tataacgtgtt cttaggttgt 540

tcgtctgttg cagggttaa

559

BUDAPEST TREATY ON THE INTERNATIONAL
RECOGNITION OF THE DEPOSIT OF MICROORGANISMS
FOR THE PURPOSES OF PATENT PROCEDURE

INTERNATIONAL FORM

Degussa-Hüls AG
Kantstr. 2
33790 Halle

RECEIPT IN THE CASE OF AN ORIGINAL DEPOSIT
issued pursuant to Rule 7.1 by the
INTERNATIONAL DEPOSITORY AUTHORITY
identified at the bottom of this page

I. IDENTIFICATION OF THE MICROORGANISM	
Identification reference given by the DEPOSITOR: DH5 α /pMAK705	Accession number given by the INTERNATIONAL DEPOSITORY AUTHORITY: DSM 13720
II. SCIENTIFIC DESCRIPTION AND/OR PROPOSED TAXONOMIC DESIGNATION	
<p>The microorganism identified under I. above was accompanied by:</p> <p><input checked="" type="checkbox"/> a scientific description <input checked="" type="checkbox"/> a proposed taxonomic designation</p> <p>(Mark with a cross where applicable).</p>	
III. RECEIPT AND ACCEPTANCE	
<p>This International Depository Authority accepts the microorganism identified under I. above, which was received by it on 2000-09-08 (Date of the original deposit)¹.</p>	
IV. RECEIPT OF REQUEST FOR CONVERSION	
<p>The microorganism identified under I. above was received by this International Depository Authority on (date of original deposit) and a request to convert the original deposit to a deposit under the Budapest Treaty was received by it on (date of receipt of request for conversion).</p>	
V. INTERNATIONAL DEPOSITORY AUTHORITY	
<p>Name: DSMZ-DEUTSCHE SAMMLUNG VON MIKROORGANISMEN UND ZELLKULTUREN GmbH</p> <p>Address: Mascheroder Weg 1b D-38124 Braunschweig</p>	<p>Signature(s) of person(s) having the power to represent the International Depository Authority or of authorized official(s):</p> <p><i>Dagmar Tröse</i></p> <p>Date: 2000-09-12</p>

¹ Where Rule 6.4 (d) applies, such date is the date on which the status of international depositary authority was acquired.

BUDAPEST TREATY ON THE INTERNATIONAL
RECOGNITION OF THE DEPOSIT OF MICROORGANISMS
FOR THE PURPOSES OF PATENT PROCEDURE

INTERNATIONAL FORM

Degussa-Hüls AG
Kantstr. 2
33790 Halle

VIABILITY STATEMENT
issued pursuant to Rule 10.2 by the
INTERNATIONAL DEPOSITORY AUTHORITY
identified at the bottom of this page

I. DEPOSITOR		II. IDENTIFICATION OF THE MICROORGANISM
Name: Degussa-Hüls AG Kantstr. 2 Address: 33790 Halle		Accession number given by the INTERNATIONAL DEPOSITORY AUTHORITY: DSM 13720 Date of the deposit or the transfer: 2000-09-08
III. VIABILITY STATEMENT		
The viability of the microorganism identified under II above was tested on 2000-09-08. On that date, the said microorganism was <input checked="" type="checkbox"/> viable <input type="checkbox"/> no longer viable		
IV. CONDITIONS UNDER WHICH THE VIABILITY TEST HAS BEEN PERFORMED ¹		
V. INTERNATIONAL DEPOSITORY AUTHORITY		
Name: DSMZ-DEUTSCHE SAMMLUNG VON MIKROORGANISMEN UND ZELLKULTUREN GmbH Address: Mascheroder Weg 1b D-38124 Braunschweig		Signature(s) of person(s) having the power to represent the International Depository Authority or of authorized official(s):  Date: 2000-09-12

¹ Indicate the date of original deposit or, where a new deposit or a transfer has been made, the most recent relevant date (date of the new deposit or date of the transfer).

² In the cases referred to in Rule 10.2(a) (ii) and (iii), refer to the most recent viability test.

³ Mark with a cross the applicable box.

⁴ Fill in if the information has been requested and if the results of the test were negative.

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization
International Bureau



(43) International Publication Date
17 October 2002 (17.10.2002)

PCT

(10) International Publication Number
WO 02/081721 A3

(51) International Patent Classification⁷: C12P 13/08,
13/04 // (C12P 13/08, C12R 1:19)

GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW,
MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG,
SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN,
YU, ZA, ZM, ZW.

(21) International Application Number: PCT/EP02/02419

(84) Designated States (regional): ARIPO patent (GH, GM,
KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW),
Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM),
European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR,
GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent
(BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR,
NE, SN, TD, TG).

(22) International Filing Date: 6 March 2002 (06.03.2002)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
101 16 518.8 3 April 2001 (03.04.2001) DE

Published:

- with international search report
- with (an) indication(s) in relation to deposited biological material furnished under Rule 13bis separately from the description

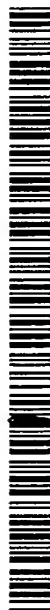
(71) Applicant: DEGUSSA AG [DE/DE]; Bennigsenplatz 1,
40474 Düsseldorf (DE).

(88) Date of publication of the international search report:
30 October 2003

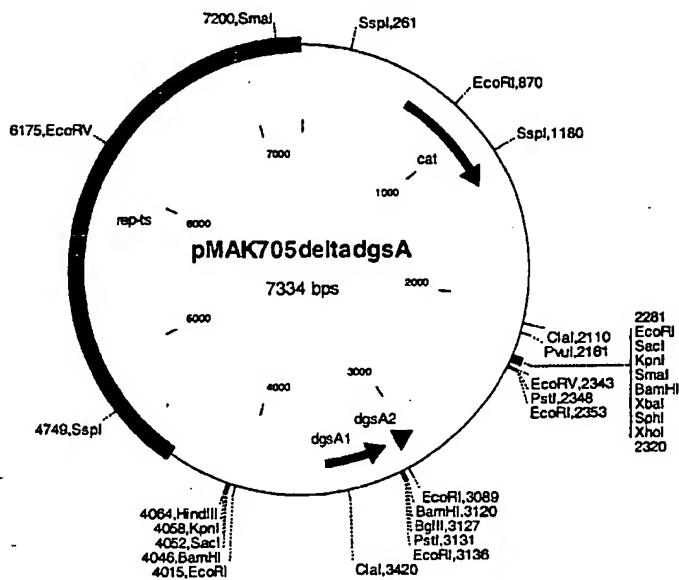
(72) Inventors: RIEPING, Mechthild; Mönckebergstrasse 1,
33619 Bielefeld (DE). HERMANN, Thomas; Zirkon-
strasse 8, 33739 Bielefeld (DE).

For two-letter codes and other abbreviations, refer to the "Guid-
ance Notes on Codes and Abbreviations" appearing at the begin-
ning of each regular issue of the PCT Gazette.

(54) Title: PROCESS FOR THE PRODUCTION OF L-AMINO ACIDS USING STRAINS OF THE FAMILY ENTEROBACTE-
RIACEAE THAT CONTAIN AN ATTENUATED DGSA GENE



WO 02/081721 A3



(57) Abstract: The invention relates to a process for the production of L-amino acids, in particular L-threonine, in which the following steps are carried out: a) fermentation of the microorganisms of the family Enterobacteriaceae producing the desired L-amino acid, in which the dgsA gene or nucleotide sequences coding therefor are attenuated, in particular are switched off, b) enrichment of the L-amino acid in the medium or in the cells of the bacteria, and c) isolation of the L-amino acid.

INTERNATIONAL SEARCH REPORT

Int'l Application No
PCT/EP 02/02419

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C12P13/08 C12P13/04 // (C12P13/08, C12R1:19)

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 7 C12P C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, PAJ, BIOSIS, MEDLINE, EMBASE

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 4 278 765 A (DEBABOV VLADIMIR G ET AL) 14 July 1981 (1981-07-14) cited in the application the whole document ----	1-7
A	MORRIS P W ET AL.: "Cloning and location of the dgsA gene of Escherichia coli." JOURNAL OF BACTERIOLOGY, vol. 163, no. 2, August 1985 (1985-08), pages 785-786, XP008016939 ISSN: 0021-9193 cited in the application the whole document ----	1-7 -/-

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

* Special categories of cited documents :

- *A* document defining the general state of the art which is not considered to be of particular relevance
- *E* earlier document but published on or after the international filing date
- *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- *O* document referring to an oral disclosure, use, exhibition or other means
- *P* document published prior to the international filing date but later than the priority date claimed

T later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

X document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

Y document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.

& document member of the same patent family

Date of the actual completion of the International search

Date of mailing of the international search report

20 May 2003

02/06/2003

Name and mailing address of the ISA
European Patent Office, P.O. 5818 Patentlaan 2
NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,
Fax (+31-70) 340-3016

Authorized officer

van de Kamp, M

INTERNATIONAL SEARCH REPORT

Int'l Application No
PCT/EP 02/02419

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	<p>HOSONO K ET AL.: "Decreasing accumulation of acetate in a rich medium by <i>Escherichia coli</i> on introduction of genes on a multicopy plasmid." <i>BIOSCIENCE, BIOTECHNOLOGY, AND BIOCHEMISTRY</i>, vol. 59, no. 2, February 1995 (1995-02), pages 256-261, XP001148978 ISSN: 0916-8451 cited in the application abstract page 260, left-hand column, line 38 -page 261, left-hand column, line 36</p> <p>---</p>	1-7
A	<p>WO 99 53035 A (ALTMAN ELLIOT ;GOKARN RAVI R (US); EITEMAN MARK A (US); UNIV GEORG) 21 October 1999 (1999-10-21) page 5, line 20-24 examples 4,7,9,10 claims 41,49 figures 1,4</p> <p>---</p>	1-7
A	<p>EP 0 643 135 A (AJINOMOTO KK) 15 March 1995 (1995-03-15) the whole document</p> <p>---</p>	1-7
A	<p>EP 0 237 819 A (KYOWA HAKKO KOGYO KK) 23 September 1987 (1987-09-23) the whole document</p> <p>---</p>	1-7
A	<p>DATABASE WPI Section Ch, Week 199148 Derwent Publications Ltd., London, GB; Class B05, AN 1991-351136 XP002241222 & JP 03 236786 A (KYOWA HAKKO KOGYO KK), 22 October 1991 (1991-10-22) abstract</p> <p>---</p>	1-7
A	<p>MICHAL G: "Biochemical pathways: an atlas of biochemistry and molecular biology" 1999 , JOHN WILEY & SONS INC. AND SPEKTRUM AKADEMISCHER VERLAG , NEW YORK - HEIDELBERG XP002240848 ISBN: 0-471-33130-9 figures 4.2-1, 4.5-1 and 4.5-2 paragraph '4.5.3!'</p> <p>---</p> <p>-/--</p>	1-7

INTERNATIONAL SEARCH REPORT

Int'l	Application No
PCT/EP 02/02419	

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	JETTEN M S M ET AL.: "Recent advances in the physiology and genetics of amino acid-producing bacteria." CRC CRITICAL REVIEWS IN BIOTECHNOLOGY, vol. 15, no. 1, 1995, pages 73-103, XP000613291 ISSN: 0738-8551 page 82, left-hand column, line 31-43 page 90, left-hand column, line 1 -page 92, left-hand column, line 17 ---	1-7
A	KRAEMER R: "Genetic and physiological approaches for the production of amino acids" JOURNAL OF BIOTECHNOLOGY, vol. 45, no. 1, 1996, pages 1-21, XP002178648 ISSN: 0168-1656 the whole document ---	1-7
A	SAWERS G: "The anaerobic degradation of L-serine and L-threonine in enterobacteria: networks of pathways and regulatory signals" ARCHIVES OF MICROBIOLOGY, vol. 171, no. 1, 1998, pages 1-5, XP002953871 ISSN: 0302-8933 the whole document ---	1-7
E	WO 02 081698 A (DEGUSSA) 17 October 2002 (2002-10-17) the whole document page 9, line 21 -page 10, line 22 claim 7 ---	1-7
E	WO 02 081722 A (DEGUSSA) 17 October 2002 (2002-10-17) the whole document page 9, line 21 -page 10, line 22 claim 7 -----	1-7

INTERNATIONAL SEARCH REPORT

Information on patent family members

Interg. Application No.

PCT/EP 02/02419

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
US 4278765	A 14-07-1981	SU HU	875663 A1 190999 B	15-09-1982 28-12-1986
WO 9953035	A 21-10-1999	AU BR CA EP JP WO US US	3555999 A 9909615 A 2325598 A1 1073722 A1 2002511250 T 9953035 A1 2003087381 A1 6455284 B1	01-11-1999 12-12-2000 21-10-1999 07-02-2001 16-04-2002 21-10-1999 08-05-2003 24-09-2002
EP 0643135	A 15-03-1995	AT CZ DE DE DK EP JP SK US EP ES WO RU	203769 T 9401658 A3 69330518 D1 69330518 T2 643135 T3 0643135 A1 3331472 B2 81994 A3 5661012 A 1020526 A2 2158867 T3 9411517 A1 2113484 C1	15-08-2001 15-12-1994 06-09-2001 08-05-2002 15-10-2001 15-03-1995 07-10-2002 10-05-1995 26-08-1997 19-07-2000 16-09-2001 26-05-1994 20-06-1998
EP 0237819	A 23-09-1987	DE DE EP JP JP KR US	3788583 D1 3788583 T2 0237819 A2 2574786 B2 63273487 A 9108634 B1 5017483 A	10-02-1994 19-05-1994 23-09-1987 22-01-1997 10-11-1988 19-10-1991 21-05-1991
JP 3236786	A 22-10-1991	JP	2877414 B2	31-03-1999
WO 02081698	A 17-10-2002	DE WO WO WO US US	10116518 A1 02081721 A2 02081698 A2 02081722 A2 2003054503 A1 2003059903 A1	17-10-2002 17-10-2002 17-10-2002 17-10-2002 20-03-2003 27-03-2003
WO 02081722	A 17-10-2002	DE WO WO WO US US	10116518 A1 02081721 A2 02081698 A2 02081722 A2 2003054503 A1 2003059903 A1	17-10-2002 17-10-2002 17-10-2002 17-10-2002 20-03-2003 27-03-2003